

Package ‘provenance’

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Title Statistical Toolbox for Sedimentary Provenance Analysis

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Description Bundles a number of established statistical methods to facilitate the visual interpretation of large datasets in sedimentary geology. Includes functionality for adaptive kernel density estimation, principal component analysis, correspondence analysis, multidimensional scaling, generalised procrustes analysis and individual differences scaling using a variety of dissimilarity measures. Univariate provenance proxies, such as single-grain ages or (isotopic) compositions are compared with the Kolmogorov-Smirnov, Kuiper, Wasserstein-2 or Sircombe-Hazelton L2 distances. Categorical provenance proxies such as chemical compositions are compared with the Aitchison and Bray-Curtis distances, and count data with the chi-square distance. Varietal data can either be converted to one or more distributional datasets, or directly compared using the multivariate Wasserstein distance. Also included are tools to plot compositional and count data on ternary diagrams and point-counting data on radial plots, to calculate the sample size required for specified levels of statistical precision, and to assess the effects of hydraulic sorting on detrital compositions. Includes an intuitive query-based user interface for users who are not proficient in R.

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ALR	<i>Additive logratio transformation</i>
-----	---

Description

Calculates Aitchison’s additive logratio transformation for a dataset of class `compositional` or a compositional data matrix.

Usage

```
ALR(x, ...)

## Default S3 method:
ALR(x, inverse = FALSE, ...)

## S3 method for class 'compositional'
ALR(x, ...)
```

Arguments

<code>x</code>	an object of class <code>compositional</code> OR a matrix of numerical values
<code>...</code>	optional arguments
<code>inverse</code>	perform the inverse inverse logratio transformation?

Value

a matrix of ALR coordinates OR an object of class `compositional` (if `inverse=TRUE`).

Examples

```
# logratio plot of trace element concentrations:
data(Namib)
alr <- ALR(Namib$Trace)
pairs(alr[,1:5])
title('log(X/Pb)')
```

amalgamate

Group components of a composition

Description

Adds several components of a composition together into a single component

Usage

```
amalgamate(x, ...)

## Default S3 method:
amalgamate(x, ...)

## S3 method for class 'compositional'
amalgamate(x, ...)

## S3 method for class 'counts'
amalgamate(x, ...)

## S3 method for class 'SRDcorrected'
amalgamate(x, ...)

## S3 method for class 'varietal'
amalgamate(x, ...)
```

Arguments

<code>x</code>	a compositional dataset
<code>...</code>	a series of new labels assigned to strings or vectors of strings denoting the components that need amalgamating

Value

an object of the same class as `X` with fewer components

Examples

```
data(Namib)
HMcomponents <- c("Zr", "tm", "rt", "TiOx", "sph", "ap", "ep",
                  "gt", "st", "amp", "cpx", "opx")
am <- amalgamate(Namib$PTHM, feldspars=c("KF", "P"),
                 lithics=c("Lm", "Lv", "Ls"), heavies=HMcomponents)
plot(ternary(am))
```

as.acomp

*create an acomp object***Description**

Convert an object of class `compositional` to an object of class `acomp` for use in the `compositions` package

Usage

```
as.acomp(x)
```

Arguments

`x` an object of class `compositional`

Value

a `data.frame`

Examples

```
data(Namib)
qfl <- ternary(Namib$PT, c('Q'), c('KF', 'P'), c('Lm', 'Lv', 'Ls'))
plot(qfl, type="QFL.dickinson")
qfl.acomp <- as.acomp(qfl)
## uncomment the next two lines to plot an error
## ellipse using the 'compositions' package:
# library(compositions)
# ellipses(mean(qfl.acomp), var(qfl.acomp), r=2)
```

as.compositional	<i>create a compositional object</i>
------------------	--------------------------------------

Description

Convert an object of class `matrix`, `data.frame` or `acomp` to an object of class `compositional`

Usage

```
as.compositional(x, method = NULL, colmap = "rainbow")
```

Arguments

<code>x</code>	an object of class <code>matrix</code> , <code>data.frame</code> or <code>acomp</code>
<code>method</code>	dissimilarity measure, either "aitchison" for Aitchison's CLR-distance or "bray" for the Bray-Curtis distance.
<code>colmap</code>	the colour map to be used in pie charts.

Value

an object of class `compositional`

Examples

```
data(Namib)
PT.acomp <- as.acomp(Namib$PT)
PT.compositional <- as.compositional(PT.acomp)
print(Namib$PT$x - PT.compositional$x)
## uncomment the following lines for an illustration of using this
## function to integrate 'provenance' with 'compositions'
# library(compositions)
# data(Glacial)
# a.glac <- acomp(Glacial)
# c.glac <- as.compositional(a.glac)
# summaryplot(c.glac, ncol=8)
```

as.counts	<i>create a counts object</i>
-----------	-------------------------------

Description

Convert an object of class `matrix` or `data.frame` to an object of class `counts`

Usage

```
as.counts(x, method = "chisq", colmap = "rainbow")
```

Arguments

x	an object of class <code>matrix</code> or <code>data.frame</code>
method	either "chisq" (for the chi-square distance) or "bray" (for the Bray-Curtis distance)
colmap	the colour map to be used in pie charts.

Value

an object of class `counts`

Examples

```
X <- matrix(c(0,100,0,30,11,2,94,36,0),nrow=3,ncol=3)
rownames(X) <- 1:3
colnames(X) <- c('a','b','c')
comp <- as.counts(X)
d <- diss(comp)
```

as.data.frame	<i>create a data.frame object</i>
---------------	-----------------------------------

Description

Convert an object of class `compositional` to a `data.frame` for use in the `robCompositions` package

Usage

```
## S3 method for class 'compositional'
as.data.frame(x, ...)
```

```
## S3 method for class 'counts'
as.data.frame(x, ...)
```

Arguments

x	an object of class <code>compositional</code>
...	optional arguments to be passed on to the generic function

Value

a `data.frame`

Examples

```
data(Namib)
Major.frame <- as.data.frame(Namib$Major)
## uncomment the next two lines to plot an error
## ellipse using the robCompositions package:
# library(robCompositions)
# plot(pcaCoDa(Major.frame))
```

as.varietal	<i>create a varietal object</i>
-------------	---------------------------------

Description

Convert an object of class `matrix` or `data.frame` to an object of class `varietal`

Usage

```
as.varietal(x, snames = NULL, method = "KS")
```

Arguments

<code>x</code>	an object of class <code>matrix</code> or <code>data.frame</code>
<code>snames</code>	either a vector of sample names, an integer marking the length of the sample name prefix, or <code>NULL</code> . <code>read.varietal</code> assumes that the row names of the <code>.csv</code> file consist of character strings marking the sample names, followed by a number.
<code>method</code>	either <code>'KS'</code> (for the Kolmogorov-Smirnov statistic) or <code>'W2'</code> (for the Wasserstein-2 distance).

Value

an object of class `varietal`

Examples

```
fn <- system.file("SNSM/Ttn_chem.csv", package="provenance")
ap1 <- read.csv(fn)
ap2 <- as.varietal(x=ap1, snames=3)
```

btev	<i>Compute the optimal kernel bandwidth</i>
------	---

Description

Uses the diffusion algorithm of Botev (2011) to calculate the bandwidth for kernel density estimation

Usage

```
btev(x)
```

Arguments

x a vector of ordinal data

Value

a scalar value with the optimal bandwidth

Author(s)

Zdravko Botev

References

Botev, Z. I., J. F. Grotowski, and D. P. Kroese. "Kernel density estimation via diffusion." *The Annals of Statistics* 38.5 (2010): 2916-2957.

Examples

```
fname <- system.file("Namib/DZ.csv", package="provenance")
bw <- btev(read.distributional(fname)$x$N1)
print(bw)
```

bray.diss	<i>Bray-Curtis dissimilarity</i>
-----------	----------------------------------

Description

Calculates the Bray-Curtis dissimilarity between two samples

Usage

```
bray.diss(x, ...)

## Default S3 method:
bray.diss(x, y, ...)

## S3 method for class 'compositional'
bray.diss(x, ...)
```

Arguments

x a vector containing the first compositional sample
 ... optional arguments
 y a vector of length(x) containing the second compositional sample

Value

a scalar value

Examples

```
data(Namib)
print(bray.diss(Namib$HM$x["N1",], Namib$HM$x["N2",]))
```

CA

Correspondence Analysis

Description

Performs Correspondence Analysis of point-counting data

Usage

```
CA(x, nf = 2, ...)
```

Arguments

x an object of class counts
 nf number of correspondence factors (dimensions)
 ... optional arguments to the corresp function of the MASS package

Value

an object of classes CA, which is synonymous to the MASS package's correspondence class.

Examples

```
data(Namib)
plot(CA(Namib$PT))
```

central.counts	<i>Calculate central compositions</i>
----------------	---------------------------------------

Description

Computes the logratio mean composition of a continuous mixture of point-counting data.

Usage

```
## S3 method for class 'counts'
central(x, ...)
```

Arguments

x	an object of class counts
...	optional arguments

Details

The central composition assumes that the observed point-counting distribution is the combination of two sources of scatter: counting uncertainty and true geological dispersion.

Value

an [5 x n] matrix with n being the number of categories and the rows containing:

theta the ‘central’ composition.

err the standard error for the central composition.

sigma the overdispersion parameter, i.e. the coefficient of variation of the underlying logistic normal distribution. `central` computes a continuous mixture model for each component (column) separately. Covariance terms are not reported.

LL the lower limit of a ‘1 sigma’ region for theta.

UL the upper limit of a ‘1 sigma’ region for theta.

mswd the mean square of the weighted deviates, a.k.a. reduced chi-square statistic.

p.value the p-value for age homogeneity

CLR

*Centred logratio transformation***Description**

Calculates Aitchison's centered logratio transformation for a dataset of class `compositional` or a compositional data matrix.

Usage

```
CLR(x, ...)
```

```
## Default S3 method:
```

```
CLR(x, inverse = FALSE, ...)
```

```
## S3 method for class 'compositional'
```

```
CLR(x, ...)
```

Arguments

`x` an object of class `compositional` OR a matrix of numerical values

`...` optional arguments

`inverse` perform the inverse inverse logratio transformation?

Value

a matrix of CLR coordinates OR an object of class `compositional` (if `inverse=TRUE`)

Examples

```
# The following code shows that applying provenance's PCA function
# to compositional data is equivalent to applying R's built-in
# princomp function to the CLR transformed data.
data(Namib)
plot(PCA(Namib$Major))
dev.new()
clrdat <- CLR(Namib$Major)
biplot(princomp(clrdat))
```

combine	<i>Combine samples of distributional data</i>
---------	---

Description

Lumps all single grain analyses of several samples together under a new name

Usage

```
combine(x, ...)
```

Arguments

x	a distributional dataset
...	a series of new labels assigned to strings or vectors of strings denoting the samples that need amalgamating

Value

a distributional data object with fewer samples than x

Examples

```
data(Namib)
combined <- combine(Namib$DZ,
                    east=c('N3', 'N4', 'N5', 'N6', 'N7', 'N8', 'N9', 'N10'),
                    west=c('N1', 'N2', 'N11', 'N12', 'T8', 'T13'))
summaryplot(KDEs(combined))
```

densities	<i>A list of rock and mineral densities</i>
-----------	---

Description

List of rock and mineral densities using the following abbreviations: Q (quartz), KF (K-feldspar), P (plagioclase), F (feldspar), Lv_f (felsic/porphyritic volcanic rock fragments), Lv_m (microlithic / porphyritic / trachitic volcanic rock fragments), Lcc (calcite), Lcd (dolomite), Lp (marl), Lch (chert), Lms (argillaceous / micaceous rock fragments), Lmv (metavolcanics), Lmf (metasediments), Lmb (metabasites), Lv (volcanic rock fragments), Lc (carbonates), Ls (sedimentary rock fragments), Lm (metamorphic rock fragments), Lu (serpentinite), mica, opaques, FeOx (Fe-oxides), turbids, zr (zircon), tm (tourmaline), rt (rutile), TiOx (Ti-oxides), sph (titanite), ap (apatite), mon (monazite), oth (other minerals), ep (epidote), othLgM (prehnite + pumpellyite + lawsonite + carpholite), gt (garnet), ctd (chloritoid), st (staurolite), and (andalusite), ky (kyanite), sil (sillimanite), amp (amphibole), px (pyroxene), cpx (clinopyroxene), opx (orthopyroxene), ol (olivine), spinel and othHM (other heavy minerals).

Author(s)

Alberto Resentini and Pieter Vermeesch

References

Resentini, A, Malusa M G and Garzanti, E. "MinSORTING: An Excel worksheet for modelling mineral grain-size distribution in sediments, with application to detrital geochronology and provenance studies." *Computers & Geosciences* 59 (2013): 90-97.

Garzanti, E, Ando, S and Vezzoli, G. "Settling equivalence of detrital minerals and grain-size dependence of sediment composition." *Earth and Planetary Science Letters* 273.1 (2008): 138-151.

See Also

restore, minsorting

Examples

```
N8 <- subset(Namib$HM,select="N8")
distribution <- minsorting(N8,densities,phi=2,sigmaphi=1,medium="air",by=0.05)
plot(distribution)
```

diss.distributional	<i>Calculate the dissimilarity matrix between two datasets of class distributional, compositional, counts or varietal</i>
---------------------	---

Description

Calculate the dissimilarity matrix between two datasets of class distributional or compositional using the Kolmogorov-Smirnov, Sircombe-Hazelton, Aitchison or Bray-Curtis distance

Usage

```
## S3 method for class 'distributional'
diss(x, method = NULL, log = FALSE, verbose = FALSE, ...)

## S3 method for class 'compositional'
diss(x, method = NULL, ...)

## S3 method for class 'counts'
diss(x, method = NULL, ...)

## S3 method for class 'varietal'
diss(x, method = NULL, ...)
```

Arguments

x	an object of class <code>distributional</code> , <code>compositional</code> or <code>counts</code>
method	if x has class <code>distributional</code> : either "KS", "Wasserstein", "Kuiper" or "SH"; if x has class <code>compositional</code> : either "aitchison" or "bray"; if x has class <code>counts</code> : either "chisq" or "bray"; if x has class <code>varietal</code> : either "KS", "W2_1D" or "W2".
log	logical. If TRUE, subjects the <code>distributional</code> data to a logarithmic transformation before calculating the Wasserstein distance.
verbose	logical. If TRUE, gives progress updates during the construction of the dissimilarity matrix.
...	optional arguments

Details

"KS" stands for the Kolmogorov-Smirnov statistic, "W2_1D" for the 1-dimensional Wasserstein-2 distance, "Kuiper" for the Kuiper statistic, "SH" for the Sircombe-Hazelton distance, "aitchison" for the Aitchison logratio distance, "bray" for the Bray-Curtis distance, "chisq" for the Chi-square distance, and "W2" for the 2-dimensional Wasserstein-2 distance.

Value

an object of class `diss`

See Also

KS.diss bray.diss SH.diss Wasserstein.diss Kuiper.diss

Examples

```
data(Namib)
print(round(100*diss(Namib$DZ)))
```

endmembers	<i>Petrographic end-member compositions</i>
------------	---

Description

A compositional dataset comprising the mineralogical compositions of the following end-members: `undissected_magmatic_arc`, `dissected_magmatic_arc`, `ophiolite`, `recycled_clastic`, `undissected_continental_block`, `transitional_continental_block`, `dissected_continental_block`, `subcreted_axial_belt` and `subducted_axial_belt`

Author(s)

Alberto Resentini and Pieter Vermeesch

References

Resentini, A, Malusa M G and Garzanti, E. "MinSORTING: An Excel worksheet for modelling mineral grain-size distribution in sediments, with application to detrital geochronology and provenance studies." *Computers & Geosciences* 59 (2013): 90-97.

Garzanti, E, Ando, S and Vezzoli, G. "Settling equivalence of detrital minerals and grain-size dependence of sediment composition." *Earth and Planetary Science Letters* 273.1 (2008): 138-151.

See Also

minsorting

Examples

```
ophiolite <- subset(endmembers,select="ophiolite")
plot(minsorting(ophiolite,densities,by=0.05))
```

get.f

Calculate the largest fraction that is likely to be missed

Description

For a given sample size, returns the largest fraction which has been sampled with $(1-p) \times 100$ % likelihood.

Usage

```
get.f(n, p = 0.05)
```

Arguments

n	the number of grains in the detrital sample
p	the required level of confidence

Value

the largest fraction that is sampled with at least $(1-p) \times 100\%$ certainty

References

Vermeesch, Pieter. "How many grains are needed for a provenance study?" *Earth and Planetary Science Letters* 224.3 (2004): 441-451.

Examples

```
print(get.f(60))
print(get.f(117))
```

get.n	<i>Calculate the number of grains required to achieve a desired level of sampling resolution</i>
-------	--

Description

Returns the number of grains that need to be analysed to decrease the likelihood of missing any fraction greater than a given size below a given level.

Usage

```
get.n(p = 0.05, f = 0.05)
```

Arguments

p	the probability that all n grains in the sample have missed at least one fraction of size f
f	the size of the smallest resolvable fraction ($0 < f < 1$)
n	the number of grains in the sample

Value

the number of grains needed to reduce the chance of missing at least one fraction f of the total population to less than p

References

Vermeesch, Pieter. "How many grains are needed for a provenance study?." Earth and Planetary Science Letters 224.3 (2004): 441-451.

Examples

```
# number of grains required to be 99% that no fraction greater than 5% was missed:  
print(get.n(0.01))  
# number of grains required to be 90% that no fraction greater than 10% was missed:  
print(get.n(p=0.1,f=0.1))
```

<code>get.p</code>	<i>Calculate the probability of missing a given population fraction</i>
--------------------	---

Description

For a given sample size, returns the likelihood of missing any fraction greater than a given size

Usage

```
get.p(n, f = 0.05)
```

Arguments

<code>n</code>	the number of grains in the detrital sample
<code>f</code>	the size of the smallest resolvable fraction ($0 < f < 1$)

Value

the probability that all n grains in the sample have missed at least one fraction of size f

References

Vermeesch, Pieter. "How many grains are needed for a provenance study?." *Earth and Planetary Science Letters* 224.3 (2004): 441-451.

Examples

```
print(get.p(60))
print(get.p(117))
```

GPA	<i>Generalised Procrustes Analysis of configurations</i>
-----	--

Description

Given a number of (2D) configurations, this function uses a combination of transformations (reflections, rotations, translations and scaling) to find a 'consensus' configuration which best matches all the component configurations in a least-squares sense.

Usage

```
GPA(X, scale = TRUE)
```

Arguments

<code>X</code>	a list of dissimilarity matrices
<code>scale</code>	boolean flag indicating if the transformation should include the scaling operation

Value

a two column vector with the coordinates of the group configuration

See Also

procrustes

indscal

Individual Differences Scaling of provenance data

Description

Performs 3-way Multidimensional Scaling analysis using Carroll and Chang (1970)'s INdividual Differences SCALing method as implemented using De Leeuw and Mair (2011)'s stress majorization algorithm.

Usage

```
indscal(..., type = "ordinal", itmax = 1000)
```

Arguments

...	a sequence of datasets of class <code>distributional</code> , <code>compositional</code> , <code>counts</code> or <code>varietal</code> , OR a single object of class <code>varietal</code> .
type	is either "ratio" or "ordinal"
itmax	Maximum number of iterations

Value

an object of class `INDSCAL`, i.e. a list containing the following items:

`delta`: Observed dissimilarities

`obsdiss`: List of observed dissimilarities, normalized

`confdiss`: List of configuration dissimilarities

`conf`: List of matrices of final configurations

`gspace`: Joint configurations aka group stimulus space

`cweights`: Configuration weights

`stress`: Stress-1 value

`spp`: Stress per point

`sps`: Stress per subject (matrix)

`ndim`: Number of dimensions

`model`: Type of smacof model

`niter`: Number of iterations

`nobj`: Number of objects

Author(s)

Jan de Leeuw and Patrick Mair

References

de Leeuw, J., & Mair, P. (2009). Multidimensional scaling using majorization: The R package smacof. *Journal of Statistical Software*, 31(3), 1-30, <<https://www.jstatsoft.org/v31/i03/>>

Examples

```
## Not run:
attach(Namib)
plot(indscal(DZ,HM,PT,Major,Trace))

## End(Not run)
```

KDE

Create a kernel density estimate

Description

Turns a vector of numbers into an object of class KDE using a combination of the Botev (2010) bandwidth selector and the Abramson (1982) adaptive kernel bandwidth modifier.

Usage

```
KDE(x, from = NA, to = NA, bw = NA, adaptive = TRUE, log = FALSE, n = 512, ...)
```

Arguments

x	a vector of numbers
from	minimum age of the time axis. If NULL, this is set automatically
to	maximum age of the time axis. If NULL, this is set automatically
bw	the bandwidth of the KDE. If NULL, bw will be calculated automatically using <code>botev()</code>
adaptive	boolean flag controlling if the adaptive KDE modifier of Abramson (1982) is used
log	transform the ages to a log scale if TRUE
n	horizontal resolution of the density estimate
...	optional arguments to be passed on to <code>density</code>

Value

an object of class KDE, i.e. a list containing the following items:

x: horizontal plot coordinates

y: vertical plot coordinates

bw: the base bandwidth of the density estimate

ages: the data values from the input to the KDE function

See Also

KDEs

Examples

```
data(Namib)
samp <- Namib$DZ$x[['N1']]
dens <- KDE(samp,0,3000,kernel="epanechnikov")
plot(dens)
```

KDEs

Generate an object of class KDEs

Description

Convert a dataset of class distributional into an object of class KDEs for further processing by the summaryplot function.

Usage

```
KDEs(
  x,
  from = NA,
  to = NA,
  bw = NA,
  samebandwidth = TRUE,
  adaptive = TRUE,
  normalise = FALSE,
  log = FALSE,
  n = 512,
  ...
)
```

Arguments

<code>x</code>	an object of class <code>distributional</code>
<code>from</code>	minimum limit of the x-axis.
<code>to</code>	maximum limit of the x-axis.
<code>bw</code>	the bandwidth of the kernel density estimates. If <code>bw = NA</code> , the bandwidth will be set automatically using <code>botev()</code>
<code>samebandwidth</code>	boolean flag indicating whether the same bandwidth should be used for all samples. If <code>samebandwidth = TRUE</code> and <code>bw = NULL</code> , then the function will use the median bandwidth of all the samples.
<code>adaptive</code>	boolean flag switching on the adaptive bandwidth modifier of Abramson (1982)
<code>normalise</code>	boolean flag indicating whether or not the KDEs should all integrate to the same value.
<code>log</code>	boolean flag indicating whether the data should be plotted on a logarithmic scale.
<code>n</code>	horizontal resolution of the density estimates
<code>...</code>	optional parameters to be passed on to <code>density</code>

Value

an object of class `KDEs`, i.e. a list containing the following items:

`kdes`: a named list with objects of class `KDE`

`from`: the beginning of the common time scale

`to`: the end of the common time scale

`themax`: the maximum probability density of all the KDEs

`pch`: the plot symbol to be used by `plot.KDEs`

`xlabel`: the x-axis label to be used by `plot.KDEs`

See Also

`KDE`

Examples

```
data(Namib)
KDEs <- KDEs(Namib$DZ, 0, 3000, pch=NA)
summaryplot(KDEs, ncol=3)
```

KS.diss	<i>Kolmogorov-Smirnov dissimilarity</i>
---------	---

Description

Returns the Kolmogorov-Smirnov dissimilarity between two samples

Usage

```
KS.diss(x, ...)

## Default S3 method:
KS.diss(x, y, ...)

## S3 method for class 'distributional'
KS.diss(x, ...)
```

Arguments

x	the first sample as a vector
...	optional arguments
y	the second sample as a vector

Value

a scalar value representing the maximum vertical distance between the two cumulative distributions

Examples

```
data(Namib)
print(KS.diss(Namib$DZ$x[['N1']], Namib$DZ$x[['T8']]))
```

Kuiper.diss	<i>Kuiper dissimilarity</i>
-------------	-----------------------------

Description

Returns the Kuiper dissimilarity between two samples

Usage

```
Kuiper.diss(x, ...)

## Default S3 method:
Kuiper.diss(x, y, ...)

## S3 method for class 'distributional'
Kuiper.diss(x, ...)
```

Arguments

x	the first sample as a vector
...	optional arguments
y	the second sample as a vector

Value

a scalar value representing the sum of the maximum vertical distances above and below the cumulative distributions of x and y

Examples

```
data(Namib)
print(Kuiper.diss(Namib$DZ$x[['N1']], Namib$DZ$x[['T8']]))
```

lines.ternary	<i>Ternary line plotting</i>
---------------	------------------------------

Description

Add lines to an existing ternary diagram

Usage

```
## S3 method for class 'ternary'
lines(x, ...)
```

Arguments

x	an object of class ternary, or a three-column data frame or matrix
...	optional arguments to the generic lines function

Examples

```
tern <- ternary(Namib$PT, 'Q', c('KF', 'P'), c('Lm', 'Lv', 'Ls'))
plot(tern, pch=21, bg='red', labels=NULL)
middle <- matrix(c(0.01, 0.49, 0.01, 0.49, 0.98, 0.02), 2, 3)
lines(ternary(middle))
```

Description

Performs classical or nonmetric Multidimensional Scaling analysis of provenance data

Usage

```
MDS(x, ...)

## Default S3 method:
MDS(x, classical = FALSE, k = 2, ...)

## S3 method for class 'compositional'
MDS(x, classical = FALSE, k = 2, ...)

## S3 method for class 'counts'
MDS(x, classical = FALSE, k = 2, ...)

## S3 method for class 'distributional'
MDS(x, classical = FALSE, k = 2, nb = 0, ...)

## S3 method for class 'varietal'
MDS(x, classical = FALSE, k = 2, nb = 0, ...)
```

Arguments

<code>x</code>	an object of class <code>distributional</code> , <code>compositional</code> , <code>counts</code> , <code>varietal</code> or <code>diss</code>
<code>...</code>	optional arguments If <code>x</code> has class <code>distributional</code> , <code>...</code> is passed on to <code>diss.distributional</code> . If <code>x</code> has class <code>compositional</code> , <code>...</code> is passed on to <code>diss.compositional</code> . If <code>x</code> has class <code>counts</code> , <code>...</code> is passed on to <code>diss.counts</code> . If <code>x</code> has class <code>varietal</code> , <code>...</code> is passed on to <code>diss.varietal</code> . Otherwise, <code>...</code> is passed on to <code>cmdscale</code> (if <code>classical=TRUE</code>), to <code>isoMDS</code> (if <code>classical=FALSE</code>).
<code>classical</code>	boolean flag indicating whether classical (<code>TRUE</code>) or nonmetric (<code>FALSE</code>) MDS should be used
<code>k</code>	the desired dimensionality of the solution
<code>nb</code>	number of bootstrap resamples. If <code>nb>0</code> , then <code>plot.MDS(...)</code> will visualise the sampling uncertainty as polygons (inspired by Nordsvan et al. 2020). The bigger <code>nb</code> , the slower the calculations. <code>nb=10</code> seems a good compromise.

Value

an object of class MDS, i.e. a list containing the following items:

points: a two column vector of the fitted configuration

classical: a boolean flag indicating whether the MDS configuration was obtained by classical (TRUE) or nonmetric (FALSE) MDS.

diss: the dissimilarity matrix used for the MDS analysis

stress: (only if classical=TRUE) the final stress achieved (in percent)

References

Nordsvan, A.R., Kirscher, U., Kirkland, C.L., Barham, M. and Brennan, D.T., 2020. Resampling (detrital) zircon age distributions for accurate multidimensional scaling solutions. *Earth-Science Reviews*, p.103149.

Vermeesch, P., 2013, Multi-sample comparison of detrital age distributions. *Chemical Geology* v.341, 140-146, doi:10.1016/j.chemgeo.2013.01.010

Examples

```
data(Namib)
plot(MDS(Namib$Major,classical=TRUE))
```

minsorting

Assess settling equivalence of detrital components

Description

Models grain size distribution of minerals and rock fragments of different densities

Usage

```
minsorting(
  X,
  dens,
  sname = NULL,
  phi = 2,
  sigmaphi = 1,
  medium = "freshwater",
  from = -2.25,
  to = 5.5,
  by = 0.25
)
```

Arguments

<code>X</code>	an object of class <code>compositional</code>
<code>dens</code>	a vector of mineral and rock densities
<code>sname</code>	sample name if unspecified, the first sample of the dataset will be used
<code>phi</code>	the mean grain size of the sample in Krumbein's phi units
<code>sigmaphi</code>	the standard deviation of the grain size distribution, in phi units
<code>medium</code>	the transport medium, one of either "air", "freshwater" or "seawater"
<code>from</code>	the minimum grain size to be evaluated, in phi units
<code>to</code>	the maximum grain size to be evaluated, in phi units
<code>by</code>	the grain size interval of the output table, in phi units

Value

an object of class `minsorting`, i.e. a list with two tables:

`mfract`: the grain size distribution of each mineral (sum of the columns = 1)

`mcomp`: the composition of each grain size fraction (sum of the rows = 1)

Author(s)

Alberto Resentini and Pieter Vermeesch

References

Resentini, A, Malusa, M G and Garzanti, E. "MinSORTING: An Excel worksheet for modelling mineral grain-size distribution in sediments, with application to detrital geochronology and provenance studies." *Computers & Geosciences* 59 (2013): 90-97.

Garzanti, E, Ando, S and Vezzoli, G. "Settling equivalence of detrital minerals and grain-size dependence of sediment composition." *Earth and Planetary Science Letters* 273.1 (2008): 138-151.

See Also

`restore`

Examples

```
data(endmembers,densities)
distribution <- minsorting(endmembers,densities,sname='ophiolite',phi=2,
                          sigmaphi=1,medium="seawater",by=0.05)
plot(distribution,cumulative=FALSE)
```

Namib

An example dataset

Description

A large dataset of provenance data from Namibia comprised of 14 sand samples from the Namib Sand Sea and 2 samples from the Orange River.

Details

Namib is a list containing the following 6 items:

DZ: a distributional dataset containing the zircon U-Pb ages for ca. 100 grains from each sample, as well as their (1-sigma) analytical uncertainties.

PT: a compositional dataset with the bulk petrography of the samples, i.e. the quartz ('Q'), K-feldspar ('KF'), plagioclase ('P'), and lithic fragments of metamorphic ('Lm'), volcanic ('Lv') and sedimentary ('Ls') origin.

HM: a compositional dataset containing the heavy mineral composition of the samples, comprised of zircon ('zr'), tourmaline ('tm'), rutile ('rt'), Ti-oxides ('TiOx'), titanite ('sph'), apatite ('ap'), epidote ('ep'), garnet ('gt'), staurolite ('st'), andalusite ('and'), kyanite ('ky'), sillimanite ('sil'), amphibole ('amp'), clinopyroxene ('cpx') and orthopyroxene ('opx').

PTHM: a compositional dataset combining the variables contained in PT and HM plus 'mica', 'opaques', 'turbids' and 'other' transparent heavy minerals ('LgM'), normalised to 100.

Major: a compositional dataset listing the concentrations (in wt TiO₂, P₂O₅ and MnO).

Trace: a compositional data listing the concentrations (in ppm) of Rb, Sr, Ba, Sc, Y, La, Ce, Pr, Nd, Sm, Gd, Dy, Er, Yb, Th, U, Zr, Hf, V, Nb, Cr, Co, Ni, Cu, Zn, Ga and Pb.

Author(s)

Pieter Vermeesch and Eduardo Garzanti

References

Vermeesch, P. and Garzanti, E., Making geological sense of 'Big Data' in sedimentary provenance analysis, *Chemical Geology* 409 (2015) 20-27

Examples

```
samp <- Namib$DZ$x[['N1']]
dens <- KDE(samp,0,3000)
plot(dens)
```

PCA

*Principal Component Analysis***Description**

Performs PCA of compositional data using a centred logratio distance

Usage

```
PCA(x, ...)
```

Arguments

`x` an object of class `compositional`
`...` optional arguments to R's `princomp` function

Value

an object of classes `PCA`, which is synonymous to the stats package's `prcomp` class.

Examples

```
data(Namib)
plot(MDS(Namib$Major, classical=TRUE))
dev.new()
plot(PCA(Namib$Major), asp=1)
print("This example demonstrates the equivalence of classical MDS and PCA")
```

plot.CA

*Point-counting biplot***Description**

Plot the results of a correspondence analysis as a biplot

Usage

```
## S3 method for class 'CA'
plot(x, labelcol = "black", vectorcol = "red", components = c(1, 2), ...)
```

Arguments

`x` an object of class `CA`
`labelcol` colour of the sample labels (may be a vector).
`vectorcol` colour of the vector loadings for the variables
`components` two-element vector of components to be plotted
`...` optional arguments of the generic `biplot` function

See Also

CA

Examples

```
data(Namib)
plot(CA(Namib$PT))
```

plot.compositional	<i>Plot a pie chart</i>
--------------------	-------------------------

Description

Plots an object of class `compositional` as a pie chart

Usage

```
## S3 method for class 'compositional'
plot(x, sname, annotate = TRUE, colmap = NULL, ...)
```

Arguments

<code>x</code>	an object of class <code>compositional</code>
<code>sname</code>	the sample name
<code>annotate</code>	a boolean flag controlling if the pies of the pie-chart should be labelled
<code>colmap</code>	an optional string with the name of one of R's built-in colour palettes (e.g., <code>heat.colors</code> , <code>terrain.colors</code> , <code>topo.colors</code> , <code>cm.colors</code>), which are to be used for plotting the data.
<code>...</code>	optional parameters to be passed on to the graphics object

Examples

```
data(Namib)
plot(Namib$Major, 'N1', colmap='heat.colors')
```

plot.distributional *Plot continuous data as histograms or cumulative age distributions*

Description

Plot one or several samples from a distributional dataset as a histogram or Cumulative Age Distributions (CAD).

Usage

```
## S3 method for class 'distributional'
plot(
  x,
  snames = NULL,
  annotate = TRUE,
  CAD = FALSE,
  pch = NA,
  verticals = TRUE,
  colmap = NULL,
  ...
)
```

Arguments

x	an object of class distributional
snames	a string or a vector of string with the names of the samples that need plotting if snames is a vector, then the function will default to a CAD.
annotate	boolean flag indicating whether the x- and y-axis should be labelled
CAD	boolean flag indicating whether the data should be plotted as a cumulative age distribution or a histogram. For multi-sample plots, the function will override this value with TRUE.
pch	an optional symbol to mark the sample points along the CAD
verticals	boolean flag indicating if the horizontal lines of the CAD should be connected by vertical lines
colmap	an optional string with the name of one of R's built-in colour palettes (e.g., heat.colors, terrain.colors, topo.colors, cm.colors), which are to be used for plotting the data.
...	optional arguments to the generic plot function

Examples

```
data(Namib)
plot(Namib$DZ, c('N1', 'N2'))
```

plot.GPA	<i>Plot a Procrustes configuration</i>
----------	--

Description

Plots the group configuration of a Generalised Procrustes Analysis

Usage

```
## S3 method for class 'GPA'
plot(x, pch = NA, pos = NULL, col = "black", bg = "white", cex = 1, ...)
```

Arguments

x	an object of class GPA
pch	plot symbol
pos	position of the sample labels relative to the plot symbols if pch != NA
col	plot colour (may be a vector)
bg	background colour (may be a vector)
cex	relative size of plot symbols
...	optional arguments to the generic plot function

See Also

procrustes

Examples

```
data(Namib)
GPA <- procrustes(Namib$DZ, Namib$HM)
coast <- c('N1', 'N2', 'N3', 'N10', 'N11', 'N12', 'T8', 'T13')
snames <- names(Namib$DZ)
bgcol <- rep('yellow', length(snames))
bgcol[which(snames %in% coast)] <- 'red'
plot(GPA, pch=21, bg=bgcol)
```

plot.INDSCAL

*Plot an INDSCAL group configuration and source weights***Description**

Given an object of class INDSCAL, generates two plots: the group configuration and the subject weights. Together, these describe a 3-way MDS model.

Usage

```
## S3 method for class 'INDSCAL'
plot(
  x,
  asp = 1,
  pch = NA,
  pos = NULL,
  col = "black",
  bg = "white",
  cex = 1,
  xlab = "X",
  ylab = "Y",
  xaxt = "n",
  yaxt = "n",
  option = 2,
  ...
)
```

Arguments

x	an object of class INDSCAL
asp	the aspect ratio of the plot
pch	plot symbol (may be a vector)
pos	position of the sample labels relative to the plot symbols if pch != NA
col	plot colour (may be a vector)
bg	background colour (may be a vector)
cex	relative size of plot symbols
xlab	a string with the label of the x axis
ylab	a string with the label of the y axis
xaxt	if = 's', adds ticks to the x axis
yaxt	if = 's', adds ticks to the y axis
option	either: 0: only plot the group configuration, do not show the source weights 1: only show the source weights, do not plot the group configuration 2: show both the group configuration and source weights in separate windows
...	optional arguments to the generic plot function

See Also

indscal

Examples

```
data(Namib)
coast <- c('N1', 'N2', 'N3', 'N10', 'N11', 'N12', 'T8', 'T13')
snames <- names(Namib$DZ)
pch <- rep(21, length(snames))
pch[which(snames %in% coast)] <- 22
plot(indscal(Namib$DZ, Namib$HM), pch=pch)
```

plot.KDE

Plot a kernel density estimate

Description

Plots an object of class KDE

Usage

```
## S3 method for class 'KDE'
plot(x, pch = "|", xlab = "age [Ma]", ylab = "", ...)
```

Arguments

x	an object of class KDE
pch	the symbol used to show the samples. May be a vector. Set pch = NA to turn them off.
xlab	the label of the x-axis
ylab	the label of the y-axis
...	optional parameters to be passed on to the graphics object

See Also

KDE

Examples

```
data(Namib)
samp <- Namib$DZ$x[['N1']]
dens <- KDE(samp, from=0, to=3000)
plot(dens)
```

plot.KDEs	<i>Plot one or more kernel density estimates</i>
-----------	--

Description

Plots an object of class KDEs

Usage

```
## S3 method for class 'KDEs'
plot(x, sname = NA, annotate = TRUE, pch = "|", ...)
```

Arguments

x	an object of class KDEs
sname	optional sample name. If sname=NA, all samples are shown on a summary plot
annotate	add a time axis?
pch	symbol to be used to mark the sample points along the x-axis. Change to NA to omit.
...	optional parameters to be passed on to the summaryplot function

See Also

KDEs summaryplot

Examples

```
data(Namib)
kdes <- KDEs(Namib$DZ)
plot(kdes, ncol=2)
```

plot.MDS	<i>Plot an MDS configuration</i>
----------	----------------------------------

Description

Plots the coordinates of a multidimensional scaling analysis as an X-Y scatter plot or ‘map’ and, if x\$classical = FALSE, a Shepard plot.

Usage

```
## S3 method for class 'MDS'
plot(
  x,
  nnlines = FALSE,
  pch = NA,
  pos = NULL,
  cex = 1,
  col = "black",
  bg = "white",
  oma = rep(1, 4),
  mar = rep(2, 4),
  mgp = c(2, 1, 0),
  xpd = NA,
  Shepard = 2,
  ...
)
```

Arguments

<code>x</code>	an object of class MDS
<code>nnlines</code>	if TRUE, draws nearest neighbour lines
<code>pch</code>	plot character (see ?plot for details). May be a vector.
<code>pos</code>	position of the sample labels relative to the plot symbols if <code>pch != NA</code>
<code>cex</code>	relative size of plot symbols (see ?par for details)
<code>col</code>	plot colour (may be a vector)
<code>bg</code>	background colour (may be a vector)
<code>oma</code>	A vector of the form <code>c(bottom, left, top, right)</code> giving the size of the outer margins in lines of text.
<code>mar</code>	A numerical vector of the form <code>c(bottom, left, top, right)</code> that gives the number of lines of margin to be specified on the four sides of the plot.
<code>mgp</code>	The margin line (in mex units) for the axis title, axis labels and axis line. See ?par for further details.
<code>xpd</code>	A logical value or NA. See ?par for further details.
<code>Shepard</code>	either: 0: only plot the MDS configuration, do not show the Shepard plot 1: only show the Shepard plot, do not plot the MDS configuration 2: show both the MDS configuration and Shepard plot in separate windows
<code>...</code>	optional arguments to the generic plot function

See Also

MDS

Examples

```

data(Namib)
mds <- MDS(Namib$DZ)
coast <- c('N1', 'N2', 'N3', 'N10', 'N11', 'N12', 'T8', 'T13')
snames <- names(Namib$DZ)
bgcol <- rep('yellow', length(snames))
bgcol[which(snames %in% coast)] <- 'red'
plot(mds, pch=21, bg=bgcol)

```

plot.minsorting	<i>Plot inferred grain size distributions</i>
-----------------	---

Description

Plot the grain size distributions of the different minerals under consideration

Usage

```

## S3 method for class 'minsoring'
plot(x, cumulative = FALSE, components = NULL, ...)

```

Arguments

x	an object of class minsoring
cumulative	boolean flag indicating whether the grain size distribution should be plotted as a density or cumulative probability curve.
components	string or list of strings with the names of a subcomposition that needs plotting
...	optional parameters to be passed on to graphics::matplot (see ?par for details)

See Also

minsoring

Examples

```

data(endmembers, densities)
OPH <- subset(endmembers, select="ophiolite")
distribution <- minsoring(OPH, densities, phi=2, sigmaphi=1,
                          medium="air", by=0.05)
plot(distribution, components=c('F', 'px', 'opaques'))

```

`plot.PCA`*Compositional biplot*

Description

Plot the results of a principal components analysis as a biplot

Usage

```
## S3 method for class 'PCA'
plot(
  x,
  labelcol = "black",
  vectorcol = "red",
  choices = 1L:2L,
  scale = 1,
  pc.biplot = FALSE,
  ...
)
```

Arguments

<code>x</code>	an object of class PCA
<code>labelcol</code>	colour(s) of the sample labels (may be a vector).
<code>vectorcol</code>	colour of the vector loadings for the variables
<code>choices</code>	see the help pages of the generic biplot function.
<code>scale</code>	see the help pages of the generic biplot function.
<code>pc.biplot</code>	see the help pages of the generic biplot function.
<code>...</code>	optional arguments of the generic biplot function

See Also

PCA

Examples

```
data(Namib)
plot(PCA(Namib$Major))
```

plot.ternary	<i>Plot a ternary diagram</i>
--------------	-------------------------------

Description

Plots triplets of compositional data on a ternary diagram

Usage

```
## S3 method for class 'ternary'
plot(
  x,
  type = "grid",
  pch = NA,
  pos = NULL,
  labels = names(x),
  showpath = FALSE,
  bg = NA,
  col = "cornflowerblue",
  ticks = seq(0, 1, 0.25),
  ticklength = 0.02,
  lty = 2,
  lwd = 1,
  ...
)
```

Arguments

x	an object of class <code>ternary</code> , or a three-column data frame or matrix
type	adds annotations to the ternary diagram, one of either <code>empty</code> , <code>grid</code> , <code>QFL.descriptive</code> , <code>QFL.folk</code> or <code>QFL.dickinson</code>
pch	plot character, see <code>?par</code> for details (may be a vector)
pos	position of the sample labels relative to the plot symbols if <code>pch != NA</code>
labels	vector of strings to be added to the plot symbols
showpath	if x has class <code>SRDcorrected</code> , and <code>showpath==TRUE</code> , the intermediate values of the SRD correction will be plotted on the ternary diagram as well as the final composition
bg	background colour for the plot symbols (may be a vector)
col	colour to be used for the background lines (if applicable)
ticks	vector of tick values between 0 and 1
ticklength	number between 0 and 1 to mark the length of the ticks
lty	line type for the annotations (see <code>type</code>)
lwd	line thickness for the annotations
...	optional arguments to the generic <code>points</code> function

See Also

ternary

Examples

```
data(Namib)
tern <- ternary(Namib$PT,'Q',c('KF','P'),c('Lm','Lv','Ls'))
plot(tern,type='QFL.descriptive',pch=21,bg='red',labels=NULL)
```

points.ternary	<i>Ternary point plotting</i>
----------------	-------------------------------

Description

Add points to an existing ternary diagram

Usage

```
## S3 method for class 'ternary'
points(x, ...)
```

Arguments

- x an object of class ternary, or a three-column data frame or matrix
- ... optional arguments to the generic points function

Examples

```
tern <- ternary(Namib$PT,'Q',c('KF','P'),c('Lm','Lv','Ls'))
plot(tern,pch=21,bg='red',labels=NULL)
# add the geometric mean composition as a yellow square:
gmean <- ternary(exp(colMeans(log(tern$x))))
points(gmean,pch=22,bg='yellow')
```

procrustes	<i>Generalised Procrustes Analysis of provenance data</i>
------------	---

Description

Given a number of input datasets, this function performs an MDS analysis on each of these and the feeds the resulting configurations into the GPA() function.

Usage

```
procrustes(...)
```

Arguments

... a sequence of datasets of classes `distributional`, `counts`, `compositional` and `varietal` OR a single object of class `varietal`.

Value

an object of class `GPA`, i.e. a list containing the following items:
points: a two column vector with the coordinates of the group configuration
labels: a list with the sample names

Author(s)

Pieter Vermeesch

References

Gower, J.C. (1975). Generalized Procrustes analysis, *Psychometrika*, 40, 33-50.

See Also

`GPA`

Examples

```
data(Namib)
gpa1 <- procrustes(Namib$DZ, Namib$HM)
plot(gpa1)

data(SNSM)
gpa2 <- procrustes(SNSM$ap)
plot(gpa2)
```

provenance

Menu-based interface for provenance

Description

For those less familiar with the syntax of the R programming language, the `provenance()` function provides a user-friendly way to access the most important functionality in the form of a menu-based query interface. Further details and examples are provided on <https://www.ucl.ac.uk/~ucfbpve/provenance/>

`provenance` provides statistical tools to interpret large amounts of distributional (single grain analyses) and compositional (mineralogical and bulk chemical) data from the command line, or using a menu-based user interface.

Usage

```
provenance()
```

Details

A list of documented functions may be viewed by typing `help(package='provenance')`. Detailed instructions are provided at <https://www.ucl.ac.uk/~ucfbpve/provenance/> and in the Sedimentary Geology paper by Vermeesch, Resentini and Garzanti (2016).

Author(s)

Pieter Vermeesch

Maintainer: Pieter Vermeesch <p.vermeesch@ucl.ac.uk>

References

Vermeesch, P., Resentini, A. and Garzanti, E., an R package for statistical provenance analysis, Sedimentary Geology, doi:10.1016/j.sedgeo.2016.01.009.

Vermeesch, P., Resentini, A. and Garzanti, E., 2016, An R package for statistical provenance analysis, Sedimentary Geology, 336, 14-25.

See Also

Useful links:

- <https://www.ucl.ac.uk/~ucfbpve/provenance/>

radialplot.counts	<i>Visualise point-counting data on a radial plot</i>
-------------------	---

Description

Implementation of a graphical device developed by Rex Galbraith to display several estimates of the same quantity that have different standard errors.

Usage

```
## S3 method for class 'counts'
radialplot(
  x,
  num = 1,
  den = 2,
  from = NA,
  to = NA,
  t0 = NA,
  sigdig = 2,
  show.numbers = FALSE,
  pch = 21,
  levels = NA,
  clabel = "",
  bg = c("white", "red"),
```

```

    title = TRUE,
    ...
)

```

Arguments

x	an object of class counts
num	index or name of the numerator variable
den	index or name of the denominator variable
from	minimum limit of the radial scale
to	maximum limit of the radial scale
t0	central value
sigdig	the number of significant digits of the numerical values reported in the title of the graphical output.
show.numbers	boolean flag (TRUE to show sample numbers)
pch	plot character (default is a filled circle)
levels	a vector with additional values to be displayed as different background colours of the plot symbols.
clabel	label of the colour legend
bg	a vector of two background colours for the plot symbols. If levels=NA, then only the first colour is used. If levels is a vector of numbers, then bg is used to construct a colour ramp.
title	add a title to the plot?
...	additional arguments to the generic points function

Details

The radial plot (Galbraith, 1988, 1990) is a graphical device that was specifically designed to display heteroscedastic data, and is constructed as follows. Consider a set of dates $\{t_1, \dots, t_i, \dots, t_n\}$ and uncertainties $\{s[t_1], \dots, s[t_i], \dots, s[t_n]\}$. Define $z_i = z[t_i]$ to be a transformation of t_i (e.g., $z_i = \log[t_i]$), and let $s[z_i]$ be its propagated analytical uncertainty (i.e., $s[z_i] = s[t_i]/t_i$ in the case of a logarithmic transformation). Create a scatterplot of (x_i, y_i) values, where $x_i = 1/s[z_i]$ and $y_i = (z_i - z_o)/s[z_i]$, where z_o is some reference value such as the mean. The slope of a line connecting the origin of this scatterplot with any of the (x_i, y_i) s is proportional to z_i and, hence, the date t_i . These dates can be more easily visualised by drawing a radial scale at some convenient distance from the origin and annotating it with labelled ticks at the appropriate angles. While the angular position of each data point represents the date, its horizontal distance from the origin is proportional to the precision. Imprecise measurements plot on the left hand side of the radial plot, whereas precise age determinations are found further towards the right. Thus, radial plots allow the observer to assess both the magnitude and the precision of quantitative data in one glance.

References

Galbraith, R.F., 1988. Graphical display of estimates having differing standard errors. *Technometrics*, 30(3), pp.271-281.

Galbraith, R.F., 1990. The radial plot: graphical assessment of spread in ages. *International Journal of Radiation Applications and Instrumentation. Part D. Nuclear Tracks and Radiation Measurements*, 17(3), pp.207-214.

Galbraith, R.F. and Laslett, G.M., 1993. Statistical models for mixed fission track ages. *Nuclear Tracks and Radiation Measurements*, 21(4), pp.459-470.

Examples

```
data(Namib)
radialplot(Namib$PT,num='Q',den='P')
```

read.compositional	<i>Read a .csv file with compositional data</i>
--------------------	---

Description

Reads a data table containing compositional data (e.g. chemical concentrations)

Usage

```
read.compositional(
  fname,
  method = NULL,
  colmap = "rainbow",
  sep = ",",
  dec = ".",
  row.names = 1,
  header = TRUE,
  check.names = FALSE,
  ...
)
```

Arguments

fname	a string with the path to the .csv file
method	either "bray" (for the Bray-Curtis distance) or "aitchison" (for Aitchison's central logratio distance). If omitted, the function defaults to 'aitchison', unless there are zeros present in the data.
colmap	an optional string with the name of one of R's built-in colour palettes (e.g., heat.colors, terrain.colors, topo.colors, cm.colors), which are to be used for plotting the data.
sep	the field separator character. Values on each line of the file are separated by this character.
dec	the character used in the file for decimal points.

row.names	a vector of row names. This can be a vector giving the actual row names, or a single number giving the column of the which contains the row names, or character string the name of the table column containing the row names.
header	a logical value indicating whether the file contains the names of the variables as its first line.
check.names	logical. If TRUE then the names of the variables in the frame are checked to ensure that they are syntactically variable names.
...	optional arguments to the built-in read.table function

Value

an object of class `compositional`, i.e. a list with the following items:

`x`: a data frame with the samples as rows and the categories as columns

`method`: either "aitchison" (for Aitchison's centred logratio distance) or "bray" (for the Bray-Curtis distance)

`colmap`: the colour map provided by the input argument

`name`: the name of the data object, extracted from the file path

Examples

```
fname <- system.file("Namib/Major.csv", package="provenance")
Major <- read.compositional(fname)
plot(PCA(Major))
```

read.counts	<i>Read a .csv file with point-counting data</i>
-------------	--

Description

Reads a data table containing point-counting data (e.g. petrographic, heavy mineral, palaeontological or palynological data)

Usage

```
read.counts(
  fname,
  method = "chisq",
  colmap = "rainbow",
  sep = ",",
  dec = ".",
  row.names = 1,
  header = TRUE,
  check.names = FALSE,
  ...
)
```

Arguments

fname	a string with the path to the .csv file
method	either "chisq" (for the chi-square distance) or "bray" (for the Bray-Curtis distance)
colmap	an optional string with the name of one of R's built-in colour palettes (e.g., heat.colors, terrain.colors, topo.colors, cm.colors), which are to be used for plotting the data.
sep	the field separator character. Values on each line of the file are separated by this character.
dec	the character used in the file for decimal points.
row.names	a vector of row names. This can be a vector giving the actual row names, or a single number giving the column of the which contains the row names, or character string the name of the table column containing the row names.
header	a logical value indicating whether the file contains the names of the variables as its first line.
check.names	logical. If TRUE then the names of the variables in the frame are checked to ensure that they are syntactically variable names.
...	optional arguments to the built-in read.table function

Value

an object of class counts, i.e. a list with the following items:

x: a data frame with the samples as rows and the categories as columns

colmap: the colour map provided by the input argument

name: the name of the data object, extracted from the file path

Examples

```
fname <- system.file("Namib/HM.csv", package="provenance")
Major <- read.counts(fname)
#plot(PCA(HM))
```

read.densities	<i>Read a .csv file with mineral and rock densities</i>
----------------	---

Description

Reads a data table containing densities to be used for hydraulic sorting corrections (minsorting and srd functions)

Usage

```
read.densities(
  fname,
  sep = ",",
  dec = ".",
  header = TRUE,
  check.names = FALSE,
  ...
)
```

Arguments

fname	a string with the path to the .csv file
sep	the field separator character. Values on each line of the file are separated by this character.
dec	the character used in the file for decimal points.
header	a logical value indicating whether the file contains the names of the variables as its first line.
check.names	logical. If TRUE then the names of the variables in the frame are checked to ensure that they are syntactically variable names.
...	optional arguments to the built-in read.table function

Value

a vector with mineral and rock densities

Examples

```
data(Namib,densities)
N8 <- subset(Namib$HM,select="N8")
distribution <- minsorting(N8,densities,phi=2,sigmaphi=1,medium="air",by=0.05)
plot(distribution)
```

read.distributional	<i>Read a .csv file with distributional data</i>
---------------------	--

Description

Reads a data table containing distributional data, i.e. lists of continuous data such as detrital zircon U-Pb ages.

Usage

```
read.distributional(
  fname,
  errorfile = NA,
  method = "KS",
  xlab = "age [Ma]",
  colmap = "rainbow",
  sep = ",",
  dec = ".",
  header = TRUE,
  check.names = FALSE,
  ...
)
```

Arguments

<code>fname</code>	the path of a .csv file with the input data, arranged in columns.
<code>errorfile</code>	the (optional) path of a .csv file with the standard errors of the input data, arranged by column in the same order as <code>fname</code> . Must be specified if the data are to be compared with the Sircombe-Hazelton dissimilarity.
<code>method</code>	an optional string specifying the dissimilarity measure which should be used for comparing this with other datasets. Should be one of either "KS" (for Kolmogorov-Smirnov), "Kuiper" (for Kuiper), "W2" (for "Wasserstein-2"), or "SH" (for "Sircombe and Hazelton"). If <code>method = "SH"</code> , then <code>errorfile</code> should be specified. If <code>method = "SH"</code> and <code>errorfile</code> is unspecified, then the program will default back to the Kolmogorov-Smirnov dissimilarity.
<code>xlab</code>	an optional string specifying the nature and units of the data. This string is used to label kernel density estimates.
<code>colmap</code>	an optional string with the name of one of R's built-in colour palettes (e.g., <code>heat.colors</code> , <code>terrain.colors</code> , <code>topo.colors</code> , <code>cm.colors</code>), which are to be used for plotting the data.
<code>sep</code>	the field separator character. Values on each line of the file are separated by this character.
<code>dec</code>	the character used in the file for decimal points.
<code>header</code>	a logical value indicating whether the file contains the names of the variables as its first line.
<code>check.names</code>	logical. If TRUE then the names of the variables in the frame are checked to ensure that they are syntactically variable names.
<code>...</code>	optional arguments to the built-in <code>read.csv</code> function

Value

an object of class `distributional`, i.e. a list with the following items:

`x`: a named list of vectors containing the numerical data for each sample

`err`: an (optional) named list of vectors containing the standard errors of `x`

method: either "KS" (for Kolmogorov-Smirnov), "Kuiper" (for the Kuiper statistic) or "SH" (for Sircombe Hazelton)

breaks: a vector with the locations of the histogram bin edges

xlab: a string containing the label to be given to the x-axis on all plots

colmap: the colour map provided by the input argument

name: the name of the data object, extracted from the file path

Examples

```
agefile <- system.file("Namib/DZ.csv", package="provenance")
errfile <- system.file("Namib/DZerr.csv", package="provenance")
DZ <- read.distributional(agefile, errfile)
plot(KDE(DZ$x$N1))
```

read.varietal	<i>Read a .csv file with varietal data</i>
---------------	--

Description

Reads a data table containing compositional data (e.g. chemical concentrations) for multiple grains and multiple samples

Usage

```
read.varietal(
  fname,
  snames = NULL,
  sep = ",",
  dec = ".",
  method = "KS",
  check.names = FALSE,
  row.names = 1,
  ...
)
```

Arguments

fname	file name (character string)
snames	either a vector of sample names, an integer marking the length of the sample name prefix, or NULL. read.varietal assumes that the row names of the .csv file consist of character strings marking the sample names, followed by a number.
sep	the field separator character. Values on each line of the file are separated by this character.
dec	the character used in the file for decimal points.

method	an optional string specifying the dissimilarity measure which should be used for comparing this with other datasets. Should be one of either "KS" (for Kolmogorov-Smirnov) or "Kuiper" (for Kuiper)
check.names	logical. If TRUE then the names of the variables in the frame are checked to ensure that they are syntactically variable names.
row.names	logical. See the documentation for the read.table function.
...	optional arguments to the built-in read.csv function

Value

an object of class `varietal`, i.e. a list with the following items:

`x`: a compositional data table

`snames`: a vector of strings corresponding to the sample names

`name`: the name of the dataset, extracted from the file path

Examples

```
fn <- system.file("SNSM/Ttn_chem.csv", package="provenance")
Ttn <- read.varietal(fname=fn, snames=3)
plot(MDS(Ttn))
```

restore	<i>Undo the effect of hydraulic sorting</i>
---------	---

Description

Restore the detrital composition back to a specified source rock density (SRD)

Usage

```
restore(X, dens, target = 2.71)
```

Arguments

X	an object of class <code>compositional</code>
dens	a vector of rock and mineral densities
target	the target density (in g/cm3)

Value

an object of class `SRDcorrected`, i.e. an object of class `compositional` which is a daughter of class `compositional` containing the restored composition, plus one additional member called `restoration`, containing the intermediate steps of the SRD correction algorithm.

Author(s)

Alberto Resentini and Pieter Vermeesch

References

Garzanti E, Ando, S and Vezzoli, G. "Settling equivalence of detrital minerals and grain-size dependence of sediment composition." *Earth and Planetary Science Letters* 273.1 (2008): 138-151.

See Also

minsorting

Examples

```
data(Namib,densities)
rescomp <- restore(Namib$PTHM,densities,2.71)
HMcomp <- c("zr","tm","rt","sph","ap","ep","gt",
            "st","amp","cpx","opx")
amcomp <- amalgamate(rescomp,Plag="P",HM=HMcomp,Opq="opaques")
plot(ternary(amcomp),showpath=TRUE)
```

SH.diss

Sircombe and Hazelton distance

Description

Calculates Sircombe and Hazelton's L2 distance between the Kernel Functional Estimates (KFEs, not to be confused with Kernel Density Estimates!) of two samples with specified analytical uncertainties

Usage

```
SH.diss(x, i, j, c.con = 0)
```

Arguments

x	an object of class <code>distributional</code>
i	index of the first sample
j	index of the second sample
c.con	smoothing bandwidth of the kernel functional estimate

Value

a scalar value expressing the L2 distance between the KFEs of samples i and j

Author(s)

Keith Sircombe and Martin Hazelton

References

Sircombe, K. N., and M. L. Hazelton. "Comparison of detrital zircon age distributions by kernel functional estimation." *Sedimentary Geology* 171.1 (2004): 91-111.

See Also

KS.diss

Examples

```
datfile <- system.file("Namib/DZ.csv",package="provenance")
errfile <- system.file("Namib/DZerr.csv",package="provenance")
DZ <- read.distributional(datfile,errfile)
d <- SH.diss(DZ,1,2)
print(d)
```

SNSM

varietal data example

Description

A list of varietal datasets including detrital zircon (zr), apatite (ap) and titanite (tit) compositions from the Sierra Nevada de Santa Marta, provided by L. Caracciolo (FAU Erlangen).

Author(s)

Luca Caracciolo, Diana Hatzenbuehler and David Chew.

Examples

```
plot(MDS(SNSM$tit))
```

subset

Get a subset of provenance data

Description

Return a subset of provenance data according to some specified indices

Usage

```
## S3 method for class 'distributional'
subset(x, subset = NULL, select = NULL, ...)

## S3 method for class 'compositional'
subset(x, subset = NULL, components = NULL, select = NULL, ...)

## S3 method for class 'counts'
subset(x, subset = NULL, components = NULL, select = NULL, ...)

## S3 method for class 'varietal'
subset(x, subset = NULL, components = NULL, select = NULL, ...)
```

Arguments

x	an object of class distributional, compositional, counts or varietal.
subset	logical expression indicating elements or rows to keep: missing values are taken as false.
select	a vector of sample names
...	optional arguments for the generic subset function
components	vector of categories (column names) to keep

Value

an object of the same class as x

See Also

[amalgamate](#), [combine](#)

Examples

```
data(Namib)
coast <- c("N1", "N2", "T8", "T13", "N12", "N13")
ZTRcoast <- subset(Namib$HM, select=coast, components=c('gt', 'cpx', 'ep'))
DZcoast <- subset(Namib$DZ, select=coast)
summaryplot(ZTRcoast, KDEs(DZcoast), ncol=2)
```

summaryplot

Joint plot of several provenance datasets

Description

Arranges kernel density estimates and pie charts in a grid format

Usage

```
summaryplot(..., ncol = 1, pch = NA)
```

Arguments

... a sequence of datasets of class `compositional`, `distributional`, `counts` or `KDEs`.

ncol the number of columns

pch (optional) symbol to be used to mark the sample points along the x-axis of the KDEs (if appropriate).

Value

a summary plot of all the data comprised of KDEs for the datasets of class `KDEs`, pie charts for those of class `compositional` or `counts` and histograms for those of class `distributional`.

See Also

[KDEs](#)

Examples

```
data(Namib)
KDEs <- KDEs(Namib$DZ,0,3000)
summaryplot(KDEs,Namib$HM,Namib$PT,ncol=2)
```

ternary

Define a ternary composition

Description

Create an object of class `ternary`

Usage

```
ternary(X, x = 1, y = 2, z = 3)
```

Arguments

X an object of class `compositional` OR a matrix or data frame with numerical data

x string/number or a vector of strings/numbers indicating the variables/indices making up the first subcomposition of the ternary system.

y second (set of) variables

z third (set of) variables

Value

an object of class ternary, i.e. a list containing:

x: a three column matrix (or vector) of ternary compositions.

and (if X is of class SRDcorrected)

restoration: a list of intermediate ternary compositions inherited from the SRD correction

See Also

restore

Examples

```
data(Namib)
tern <- ternary(Namib$PT,c('Q'),c('KF','P'),c('Lm','Lv','Ls'))
plot(tern,type="QFL")
```

ternary.ellipse	<i>Ternary confidence ellipse</i>
-----------------	-----------------------------------

Description

plot a $100(1 - \alpha)\%$ confidence region around the data or around its mean.

Usage

```
ternary.ellipse(x, ...)

## Default S3 method:
ternary.ellipse(x, alpha = 0.05, population = TRUE, ...)

## S3 method for class 'compositional'
ternary.ellipse(x, alpha = 0.05, population = TRUE, ...)

## S3 method for class 'counts'
ternary.ellipse(x, alpha = 0.05, population = TRUE, ...)
```

Arguments

x	an object of class ternary
...	optional formatting arguments
alpha	cutoff level for the confidence ellipse
population	show the standard deviation of the entire population or the standard error of the mean?

Examples

```
data(Namib)
tern <- ternary(Namib$Major, 'CaO', 'Na2O', 'K2O')
plot(tern)
ternary.ellipse(tern)
```

text.ternary

Ternary text plotting

Description

Add text an existing ternary diagram

Usage

```
## S3 method for class 'ternary'
text(x, labels = 1:nrow(x$x), ...)
```

Arguments

x	an object of class ternary, or a three-column data frame or matrix
labels	a character vector or expression specifying the text to be written
...	optional arguments to the generic text function

Examples

```
data(Namib)
tern <- ternary(Namib$Major, 'CaO', 'Na2O', 'K2O')
plot(tern, pch=21, bg='red', labels=NULL)
# add the geometric mean composition as a text label:
gmean <- ternary(exp(colMeans(log(tern$x))))
text(gmean, labels='geometric mean')
```

varietal2distributional

Convert varietal to distributional data

Description

Convert an object of class varietal either to a list of distributional objects by breaking it up into separate elements, or to a single distributional object corresponding to the first principal component.

Usage

```
varietal2distributional(x, bycol = FALSE, plot = FALSE)
```

Arguments

<code>x</code>	an object of class <code>varietal</code> .
<code>bycol</code>	logical. If TRUE, returns a list of distributional objects (one for each element). If FALSE, returns a single distributional object (containing the PC1 scores for each sample).
<code>plot</code>	logical. If TRUE, shows the PCA biplot that is used when <code>bycol</code> is FALSE.

Examples

```
Ttn_file <- system.file("SNSM/Ttn_chem.csv", package="provenance")
Ttn <- read.varietal(fn=Ttn_file, snames=3)
varietal2distributional(Ttn, bycol=FALSE, plot=TRUE)
```

Wasserstein.diss	<i>Wasserstein distance</i>
------------------	-----------------------------

Description

Returns the Wasserstein distance between two samples

Usage

```
Wasserstein.diss(x, ...)

## Default S3 method:
Wasserstein.diss(x, y, ...)

## S3 method for class 'distributional'
Wasserstein.diss(x, log = FALSE, ...)

## S3 method for class 'varietal'
Wasserstein.diss(x, package = "transport", verbose = FALSE, ...)
```

Arguments

<code>x</code>	the first sample as a vector
<code>...</code>	optional arguments to the <code>transport::wasserstein()</code> or <code>T4transport::wasserstein()</code> functions. Warning: the latter function is very slow.
<code>y</code>	the second sample as a vector
<code>log</code>	logical. Take the lograthm of the data before calculating the distances?
<code>package</code>	the name of the package that provides the 2D Wasserstein distance. Currently, this can be either <code>'transport'</code> or <code>T4transport</code> .
<code>verbose</code>	logical. If TRUE, gives progress updates during the construction of the dissimilarity matrix.

Value

a scalar value

Author(s)

The default S3 method was written by Pieter Vermeesch, using modified code from Dominic Schuhmacher's transport package (transport1d function), as implemented in IsoplotR.

Examples

```
data(Namib)
print(Wasserstein.diss(Namib$DZ$x[['N1']], Namib$DZ$x[['T8']]))
```

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